# **BIONETICS**

MUTAGENICITY EVALUATION

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FDA 75-86 NIACINAMIDE

FINAL REPORT

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#### MUTAGENICITY EVALUATION

OF
FDA 75-86
NIACINAMIDE
FINAL REPORT

#### SUBMITTED TO

GENETIC TOXICOLOGY BRANCH
DIVISION OF TOXICOLOGY
BUREAU OF FOODS
U.S. FOOD AND DRUG ADMINISTRATION
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LBI PROJECT NO. 2672

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#### TABLE OF CONTENTS

	P	age	No.
EVALUATION	SUMMARY		1
I.	<u>OBJECTIVE</u>	•	2
II.	MATERIALS	•	2
	A. Test Compound	•	2 2 2 3 3
III.	METHODS	• •	3
	A. Toxicity  B. Plate Tests  C. Suspension Tests  D. Preparation of Tissue Homogenates and 9,000 x g  Cell Fractions  E. Data Recording and Reporting	· ·	3 4 4 5 5
IV.	RESULTS SECTION		
	A. Solubility Properties of the Test Compound B. Toxicity and Dosage Determinations for the Test		6
	Compound	• •	6 7 7
٧.	INTERPRETATION OF RESULTS AND CONCLUSIONS	1	5
VI.	EXPLANATION OF EVALUATION PROCEDURES FOR PLATE ASSAYS	1	6
VII.	EXPLANATION OF EVALUATION PROCEDURES FOR SUSPENSION ASSA	<u>YS</u> 1	8
ADDENNTY	- Tabulation of Data		A-1



#### EVALUATION SUMMARY

The test compound, FDA 75-86, Niacinamide, did not exhibit mutagenic activity in any of the assays employed in these studies.



DATE:

July, 1977

SPONSOR: U.S. Food and Drug Administration

SUBJECT: Evaluation of Test Compound: FDA 75-86, Niacinamide

#### I. OBJECTIVE

The objective of this study was to evaluate the test compound for genetic activity in microbial assays with and without the addition of mammalian metabolic activation preparations.

#### II. **MATERIALS**

Α.

Test Compound

1.

Date Received:

December 29, 1976

2.

Description:

White crystals

#### Indicator Microorganisms В.

The following strains of indicator microorganisms were used in the evaluation:

Yeast Strain:

Saccharomyces cerevisiae, strain D4

Bacteria Strains:

Salmonella typhimurium, strains TA-1535

TA-1537

TA-1538

TA-98

TA-100

#### С. Reaction Mixture

The following reaction mixture was employed in the activation tests:

#### Component

#### Final Concentration/ml

umoles

umoles

5 µmoles

100 µmoles

33 µmoles

- TPN (sodium salt)
- 2. Glucose-6-phosphate
- 3. Sodium phosphate (dibasic)
- 4. MgCl<sub>2</sub>
- 5. KC1
- 6. Homogenate fraction equivalent to 25 mg of wet tissue.



#### D. Tissue Homogenates and Supernatants

The tissue homogenates and  $9,000 \times \underline{g}$  supernatants were prepared from tissues of the following mammalian species: Mouse - ICR random bred adult males; rat - Sprague-Dawley adult males; and monkey -  $\underline{\text{Macaca mulatta}}$  adult males.

#### E. Positive Control Compounds

Table 1 lists chemicals for positive controls in the direct and activation assays.

TABLE 1

POSITIVE CONTROLS USED IN DIRECT AND ACTIVATION ASSAYS

Assay	<u>Chemical<sup>a</sup></u>	Solvent	Probable Mutagenic Specificity
Nonactivation	Methylnitrosoguanidine	Water or saline	BPSb
	Ethylmethanesulfonate	Water or saline	BPSb
	2-Nitrofluorene	Dimethylsulfoxide <sup>C</sup>	FSb
	Quinacrine mustard	Water or saline	FSb
Activation	Dimethylnitrosamine	Water or saline	BPS <sup>b</sup>
	2-Acetylaminofluorene	Dimethylsulfoxide <sup>C</sup>	FS <sup>b</sup>
	8-Aminoquinoline	Dimethylsulfoxide <sup>C</sup>	FS <sup>b</sup>
	2-Aminoanthracene	Dimethylsulfoxide <sup>C</sup>	BPS <sup>b</sup>

a Concentrations given in the Results Section

#### III. METHODS

#### A. Toxicity

The solubility, toxicity and doses for the test chemical were determined prior to screening.

The test chemical was tested for toxicity against specific indicator strains over a range of doses to determine the 50% survival dose. Bacteria were tested in phosphate buffer, pH 7.4, for one hour at 37°C on a shaker. Yeasts were tested in phosphate buffer, pH 7.4, for four hours at 30°C on a shaker. The 50% survival concentrations and the 1/4 and 1/2 50% doses calculated.

If no toxicity was obtained for the chemical with a given strain, then a maximum dose of 5% (w/v) was used.

Unless otherwise specified, the doses calculated for the tests in buffer were applied to the activation tests. The solubility of the test chemical under treatment conditions is stated in the Results Section.



BPS = base-pair substitution; FS = frameshift
Previously shown to be non-mutagenic

### B. Plate Tests (Overlay Method)

Approximately  $10^8$  cells from an overnight culture of each indicator strain were added to test tubes containing 2.0 ml of molten agar supplemented with biotin and a trace of histidine. For nonactivation tests, the three dose levels of the test compound were added to the contents of the appropriate tubes and poured over the surfaces of selective agar plates. In activation tests 0.5 ml of a 9,000 x g tissue supernatant and required cofactors (core reaction mixture) were added to the overlay tubes. Three dose levels of the test chemical were added to the appropriate tubes, which were then mixed and the contents poured over the surface of a minimal agar (selective medium) plate and allowed to solidify. The plates were incubated for 48 to 72 hours at  $37^{\circ}\text{C}$ , and scored for the number of colonies growing on each plate. The concentrations of all chemicals are given in the Results Section. Positive and solvent controls using positive compounds that are active directly and those that require metabolic activation were run with each assay.

#### C. Suspension Tests

#### Nonactivation

Bacteria and yeast cultures of the indicator organisms were grown in complete broth, washed and resuspended in 0.9% saline to densities of 1  $\times$  10<sup>10</sup> cells/ml and 5  $\times$  10 $^{9}$  cells/ml, respectively. This constituted the working stock for tests of a group of test chemicals and their respective controls. Tests were conducted in plastic, 24-well tissue culture plates (Linbro). Cells plus appropriate volume(s) of the test chemical were added to the wells to give a final volume of 1.5 ml. The solvent replaced the test chemical in the negative controls. Treatment was at 30°C for four hours for yeast tests and at 37°C for one hour for bacterial tests. All flasks were shaken during treatment. Following treatment, the plates were set on ice. Aliquots of cells were removed, diluted in sterile saline (4°C) and plated on the appropriate complete media. Undiluted samples from flasks containing the bacteria were plated on minimal selective medium in reversion experiments. Samples from a  $10^{-1}$  dilution of treated cells were plated on the selected media for enumeration of gene conversion with strain D4. Bacterial plates were scored after incubation for 48 hours at 37°C. The yeast plates were incubated at 30°C for 3-5 days before scoring.

#### Activation

Bacteria and yeast cells were grown and prepared as described in the nonactivation tests. Measured amounts of the test and control chemicals plus 0.25 ml of the stock-cell suspension were added to wells of the Linbro plate containing the appropriate tissue fraction and reaction mixture. All flasks (bacteria and yeast) were incubated at 37°C with shaking. The treatment times as well as the dilutions, plating procedures and scoring of the plates were the same as described for nonactivation tests.



#### D. Preparation of Tissue Homogenates and 9,000 x g Cell Fractions

Male animals (except monkeys) sufficient to provide the necessary quantities of tissues were killed by cranial blow, decapitated and bled. Monkey tissues were obtained from freshly killed and bled male rhesus monkeys. Organs were immediately dissected from the animals using aseptic techniques and placed in ice-cold 0.15M KCl. Upon collection of the desired quantity of organs, they were washed twice with fresh KCl and completely homogenized with a motor-driven homogenizing unit at  $4^{\circ}$ C. The whole organ homogenate obtained from this step was divided into two samples. One sample was frozen at  $-80^{\circ}$ C and the other was centrifuged for 20 minutes at 9,000 x g in a refrigerated centrifuge. The supernatant from the centrifuged sample was retained and frozen at  $-80^{\circ}$ C. These two frozen samples were used for the activation studies. Protein and P-448 determinations were made for each lot of homogenate.

#### E. Data Recording and Reporting

#### 1. Plate test assays

The numbers of colonies on each plate were counted and recorded on printed forms. These raw data were entered into a computer program designed to print out all data by test. The data are presented as revertants per plate for each indicator strain employed in the assay. The positive and solvent controls are provided as reference points.

#### 2. Suspension assays

Following the specified incubation periods all population plates were scored by an automatic colony counter and the results from each plate of a set were recorded, in ink, on data processing forms. All minimal or other types of selective media plates were hand scored and the results recorded along with the respective population data. Other relevant experimental data were recorded on experimental definition forms. For bacteria strains the number of colonies recorded from either the population or selective plates represents that number in 1 ml of test suspension plated. The numbers recorded for the yeast strain D4 represent the number in 0.5 ml of test suspension plated. The data were then processed and printed from a computer program. All raw data sheets are dated and signed by the responsible technician.



- IV. RESULTS SECTION
- A. Solubility Properties of the Test Compound
- 1. Name or code designation of the test compound: FDA 75-86, Niacinamide
- Test solvent: \*Saline
- 3. Solubility of the test compound under treatment conditions: Soluble
- 4. Additional comments: White crystal
- B. <u>Toxicity and Dosage Determinations for the Test Compound</u>
- 1. Test date for toxicity determination: April 4, 1977
- 2. The 50% survival level was determined for bacteria and yeast indicator organisms by conducting survival curves with the test compound at the following concentrations:

#### Percent Concentration (w/v or v/v)

5.0 0.5 0.05

0.005

0.0005

3. Concentrations of the test compound used in the mutagenicity tests:

	Percent Concentration						
Test Doses	Bacteria	Yeast					
1/4 50% Survival	0.40	0.22					
1/2 50% Survival	0.80	0.44					
50% Survival	01.6	0.88					

<sup>\*</sup>The concentration of solvent was equal to the highest volume of test material added.



#### C. Plate Test Results

The plate test results are summarized in the following table. The values presented in this table are the number of revertants per plate.

#### D. Suspension Assay Results

The suspension test results for the test compound are summarized in the tables following the plate test summary. The values presented in these tables are the calculated mutation frequencies for each control and experimental test point. The first table of the suspension set presents the results for the nonactivation assays, and the second table through the fourth table of the suspension set presents the results for the activation assays. A listing of computer codes and abbreviations is included for reference. Tabulation of all raw data is provided in the Appendix.



#### SUMMARY\_OE\_IEST\_RESULIS

PLAIE IESIS

NAME OR CODE DESIGNATION OF THE TEST COMPOUND: 000098920

TEST DATE: MAY 18, 1977 8.

						B.E.Y.	E.B.I.	ANI	S P		P_L_A	J_E		
IES	$\mathbf{I}$		SPECIES	IISSUE	IA:	-1535_	IA:	-1537_	IA:	-1538_	IA:	-98	_14=	100
					1	2	1	2	1	2	1	2	1	2
1.	NON-ACI	LIVAIION												
	SOLVENT	CONTROL+			28	21	55	35	17	16	34	27	148	143
	POSITIV	/E CONTROL**			>1000	>1000	>1000	>1000	>1000	>1000	>1000	>1000	>1000	>1000
	TEST	1.60000 %			45	35	13	19	14	13	29	23	131	124
		0.80000 %			18	49	15	10	10	10	27	24	128	122
		0.40000 %			22	22	- 13	11	10	14	31	23	151	127
2.	ACILYA	LION												
	SOLVENT	CONTROL*	NOUSE	LIVER	30	31	22	23	19	10	37	32	555	195
			RAT	LIVER	26	37	20	18	19	17	39	40	147	182
			MONKEY	LIVER	18	15	17	31	23	21	36	37	192	133
	POSITI	VE CONTROL***	MOUSE	LIVER	502	490	260	256	874	911	>1000	>1000	624	889
			RAT	LIVER	274	374	241	149	938	732	>1000	>1000		>1000
			HONKEY	LIVER	370	215	173	160	738	901	>1000	937	>1000	>1000
	TEST	1.60000 %	MOUSE	LIVER	27	26	21	31	20	19	38	41	153	154
		0.80000 %	HOUSE	LIVER	25	17	19	19	14	18	39	36	171	165
		0.40000 %	MOUSE	LIVER	26	27	17	21	17	16	28	35	188	152
		1.60000 %	RAT	LIVER	20	28	20	22	13	11	36	32	128	135
		0.80000 %	RAT	LIVER	27	29	21	14	11	18	41	42	129	134
		0.40000 %	. RAT	LIVER	16	25	12	12	11	10	39	41	135	145
		1.60000 %	MONKEY	LIVER	14	26	26	23	21	23	44	41	149	141
		0.80000 %	MONKEY	LIVER	25	55	15	22	12	14	38	32	151	146
		0.40000 %	MONKEY	LIVER	20	25	23	24	18	10	30	33	147	142

<sup>.</sup> NON-ACTIVATION ASSAYS CONSIST OF THE CELLS PLUS THE TEST COMPOUND VEHICLE (SOLVENT). FOR ACTIVATION ASSAYS. THE OVERLAY CONTAINS THE ACTIVATION SYSTEM PLUS THE TEST COMPOUND VEHICLE.

**	T4-1535	MNNG	2	UG/PLA	ſΕ		***	TA-1535	ANT	H 100	UG/P	LATE	
	TA-1537	QM	20	UG/PLA	1E			TA-1537	7 AMQ	100	UG/P	LATE	
	TA-1538	NF	100	UG/PLA	ΓE			TA-1538	AAF	100	UG/P	LATE	
	TA-98	NF	100	UG/PLA	ΙE			TA-98	AAF	100	UG/P	LATE	
	TA-100	MNNG	2	UG/PLA	ΓE			TA-100	ANT	H 100	UG/P	LATE	
	HOTE	CONCEN	TRAT	IONS ARE	GIVEN	IN	MICROLITE	RS (UL)	OR HI	CROGRAM	S (UG)	PER	PLATE.

# LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM REPORT EXR34

#### COMPOUND FREQUENCY SUMMARY REPORT 07/22/77

### NONACTIVATION COMPOUND 000098920

TEST	nRG	TA100 HIS EX-8	TA1535 HIS EX-8	TA1537 H1S EX-8	TA1538 HIS EX-8	TA98 HIS EX-8	0000D4 ADE EX-5	0000D4 TRY EX-5		
			1							
NAN		85.71	3.58	11.59	5.46	13.83	19.91	6.89 -	CONTROLS	
NAP		900.49	685.45	235.32	163.30	71.70	109.36	78.06		<b></b>
NAI		81.48	4.55	3.35	4.38	5.47	40.14	6.41	TEST DATA	
NAZ		67.99	3.88	2.79	3.89	10.36	19.13	5.63		
NA3		68.17	3.76	3.93	2.67	9.79	27.94	11.93		

### LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM REPORT EXR34

#### COMPOUND FREQUENCY SUMMARY REPORT 07/22/77

SPECIES ICRFLO/MOUSE

COMPOUND 000098920

TEST	nRG	TA100 HIS EX-8	TA1535 HIS EX-8	TA1537 HIS EX-B	TA1538 HIS EX-8	TA98 HIS EX-8	000004 ADE EX-5	0000D4 TRY EX-5	
ACT	A+C	84.43	3.81	9.31	15.20	9.46	6.93	15.63	NEGATIVE CONTROLS
ACT	A-C	55.84	5.81	6.29	5.26	8.79	6.90	22.67	
ACT	ALI	57.41	5.02	12.74	5.19	21.48	10.38	8.14	
ACT	ALU	58.69	5.12			14.70			
ACT	PLI	182.07	162.45					91.82	POSITIVE CONTROLS
ACT		91.94			34.19			17.34	
ACT	LII								TEST COMPOUND
ACT	L12	72.96	3.91	16.55	4.12	21.03	8.91	2.67	
ACT	L13	71.03	5.95	16.82	3.62	18.53	5.30	1.48	
ACT	1.01	87.50	8.91	10.72	11.84	12.80	4.09	2.36	
ACT	LU2	23.72	4.11	18.88	6.98	8.73	4.42	2.35	
ACT	LU3	44.35	2.78	18.74	9.24	9.31	3.50	1.64	•

### LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM REPORT EXR34

#### COMPOUND FREQUENCY SUMMARY REPORT 07/22/77

SPECIES SPRDAW/RAT

COMPOUND 000098920

TEST	ORG	TA100 HIS EX-8	TA1535 HIS EX-8	TA1537 HIS EX-8	TA1538 HIS EX-8	TA98 HIS EX-8	0000D4 ADE EX-5	0000D4 TRY EX-S	
ACT	A+C	20.79	1.37	2.13	10.56	13.11	15.60	10.18	NEGATIVE CONTROLS
ACT	A-C	18.64	3.80	6.21	5.06	15.56	12.85	7.44	
ACT	ALI	83.48	2.64	7.26	10.82	36.48	15.01	10.34	
ACT	ALU		1.76					10.27	
ACT									POSITIVE CONTROLS
ACT	PLU		1.29	<del>-</del>	160.05				
ACT	LII				6.59			10.98	TEST COMPOUND
ACT	L15	37.61	2.36	5.13	14.68	51.31	17.16	12.53	•
ACT	L13	34.93	3.52	3.49	9.86	35.38	15.28	7.51	
ACT	LU1	76.38	2.00	6.07	16.81	34.62	13.40	4.92	
ACT	LU2	67.07	1.97	3.45	9.82	18.12	16.80	6.01	
ACT	LU3	84.86	2.43	3.66	13.02	29.24	15.95	9.99	

# LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM REPORT EXR34

#### COMPOUND FREQUENCY SUMMARY REPORT 07/22/77

SPECIES RHESUS/MONKEY

COMPOUND 000098920

TEST	он6	TA100 HIS EX-8	TA1535 HIS EX-8	TA1537 HIS EX-8	TA1538 HIS EX-8	TA98 HIS EX-8	0000D4 ADE EX-5	0000D4 TRY EX-5	
ACT	A+C	87.84	5.64	21.88	13.86	7.51	21.98	10.90	NEGATIVE CONTROLS
ACT	A-C	64.54	2.96	1.25	11.47	6.19	8.04	7.30	
ACT	ALI	80.09	5.54	4.98	12.37	14.55	16.25	6.21	
ACT	ALU	70.23	3.78	7.32	10.10	10.45	20.74	7.73	
ACT	PLI	284.60	56.90		154.01			54.23	POSITIVE CONTROLS
ACT	PLU	69.20					20.46		
ACT	LII	53.18						10.66	TEST COMPOUND
ACT	LI2	76.03	3.71	5.53	13.05	9.07	17.33	12.82	
ACT	L13	86.80	3.09	5.60	6.95	17.71	11.63	~ 4.11	
ACT	LU1	65.23	6.07	3.44	8.77	7.54	22.49	8.38	
ACT	LU2	81.52	2.62	2.80	8.02	11.16	5.90	3.27	
ACT	LU3	63.56	3.97	4.26	8.63	10.14	15.45	7.23	

### DATA TABLE TERMS AND ABBREVIATIONS

OR TERM		DEFINITION OR EXPLANATION						
COMPOUND	Client designated compound number appears in this column.							
TEST CODES	NAN NAP NA1 NA2, etc.	<pre>= Nonactivation: Solvent Control = Nonactivation: Positive Control = Nonactivation: Test Compound Dose 1 = Reflects the other dose level(s)</pre>						
	A+C A-C ALI ALU or A+T ACP ACT	<pre>= Negative Chemical Control for ACP = Activation: Solvent Control = Activation: Homogenate Control (Liver = Activation: Homogenate Control = Activation: Positive Control = Activation Test</pre>						
	LI LU KI TE 1,2, etc.	<ul> <li>Liver Tissue Activation Fraction</li> <li>Lung Tissue Activation Fraction</li> <li>Kidney Tissue Activation Fraction</li> <li>Testes Tissue Activation Fraction</li> <li>Dose Levels</li> </ul>						
CONCENTRATION	whole number f	ound dose levels are expressed as a followed by an exponent (negative) the appropriate units.						
	Example: 0025	-2PCT = 0.25 percent concentration						
POPU	raised to some	f viable cells in the plating sample exponent printed directly below the i.e., EP + $6 = \times 10^6$ ).						
MUT 1	from the sampl printed direct	f mutants or convertants obtained e plated raised to some exponent ly below the abbreviation (i.e., For strain D4, MUT 1 represents the convertants.						
MUT 2	Only used for of TRY+ conver	strain D4 and represents the number tants in the plated sample.						
FREQ 1	frequency time	mutation or gene conversion s the negative exponent ly below. For strain D4, FREQ l ADE+ value.						
FREQ 2	Only used for conversion fre	strain D4 and represents the TRY+ quency.						
CONTAM	Presence of co	ntamination on any plates.						

### DATA TABLE TERMS AND ABBREVIATIONS (continued)

DEFINITION OR EXPLANATION
2-Acetylaminofluorene
Dimethylsulfoxide
Dimethylnitrosamine
Ethylmethanesulfonate
Quinacrine Mustard
Nitrofluorene
2-Amino Anthracene
8-Amino Quinoline
Animal Strains
Sprague Dawley Rats
Flow ICR Random Bred Mice
Rhesus Monkey ( <u>Macaca mulatta</u> )
Dog, Mixed Breed
New Zealand White Rabbit
Microgram
Micromole
Adenine
Tryptophan



#### V. <u>INTERPRETATION OF RESULTS AND CONCLUSIONS</u>

The test compound, FDA 75-86, Niacinamide, was evaluated for genetic activity in a series of in vitro microbial assays with and without metabolic activation. The following results were obtained:

- A. <u>Salmonella typhimurium</u>
- 1. Plate tests

The results of these tests were negative.

2. Nonactivation suspension tests

The results of these tests were negative.

3. Activation suspension tests

The results of these tests were negative.

- B. Saccharomyces cerevisiae
- 1. Nonactivation suspension tests

The results of these tests were negative.

2. Activation suspension tests

The results of these tests were negative.

#### C. Conclusions

The test compound, FDA 75-86, Niacinamide, did not exhibit mutagenic activity in any of the assays employed in these studies.

Submitted by:

Dåvid J. Brusick, Ph.[

Director

Department of Molecular

Toxicology

Reviewed by:

Robert J. Weir, Ph.D.

Vice President



#### VI. <u>EXPLANATION OF EVALUATION PROCEDURES FOR PLATE ASSAYS</u>

Plate test data consist of direct revertant colony counts obtained from a set of selective agar plates seeded with populations of mutant cells suspended in a semisolid overlay. Because the test chemical and cells are incubated in the overlay for 2-3 days, and a few cell divisions occur during the incubation period, the test is semiquantitative in nature. Although these features of the assay reduce the quantitation of results, they provide certain advantages not contained in a quantitative suspension test.

- The small number of cell divisions permits potential mutagens to act on replicating DNA which is often more sensitive than non-replicating DNA.
- The combined incubation of the compound and the cells in the overlay permit constant exposure of the indicator cells for 2-3 days.

#### A. Surviving Populations

Plate test procedures do not permit exact quantitation of the number of cells surviving chemical treatment. At low concentrations of the test chemical, the surviving population on the treatment plates is essentially the same as the negative control plate. At high concentrations, the surviving population is usually reduced by some fraction. Our protocol normally employs dose levels that are selected such that the highest dose will show slight toxicity (as determined by subjective criteria) and several doses ranging down 1 to 2 logs lower.

#### B. Dose Response Phenomena

The demonstration of dose-related increases in mutant counts is an important criterion in establishing mutagenicity. Factors which may modify dose response results for a mutagen would be the selection of doses that are too low (usually mutagenicity and toxicity are related). If the highest dose is far lower than a toxic concentration, no increases may be observed over the dose range selected. Conversely, if the lowest dose employed is highly cytotoxic, the test chemical may kill any mutants that are induced and the compound will not appear to be mutagenic.

#### C. Control Tests

Positive and negative control assays are conducted with each experiment and consist of direct acting mutagens for nonactivation assays and mutagens that require metabolic biotransformation in activation assays. Negative controls consist of the test compound solvent in the overlay agar with the other essential components. The negative control plate for each strain gives a reference point to which the test data are compared. The positive control assay is conducted to demonstrate that the test systems are functional with known mutagens.



#### D. <u>Evaluation Criteria</u> for Ames Assay

Because the procedures used to evaluate the mutagenicity of the test chemical are semiquantitative, the criteria used to determine positive effects are inherently subjective and are based primarily on a historical data base. Most data sets are evaluated using the following criteria:

#### 1. Strains TA-1535, TA-1537, and TA-1538

If the solvent control value is within the normal range, a chemical that produces a positive dose response over three concentrations with the lowest increase equal to twice the solvent control value is considered to be mutagenic.

#### 2. Strains TA-98, TA-100, and D4

If the solvent control value is within the normal range, a chemical that produces a positive dose response over three concentrations with the highest increase equal to twice the solvent control value for TA-100 and two to three times the solvent control value for strains TA-98 and D4 is considered to be mutagenic. For these strains, the dose response increase should start at approximately the solvent control value.

#### Pattern

Because TA-1535 and TA-100 were both derived from the same parental strain (G-46) and because TA-1538 and TA-98 were both derived from the same parental strain (D3052), there is a built-in redundancy in the microbial assay. In general the two strains of a set respond to the same mutagen and such a pattern is sought. It is also anticipated that if a given strain, e.g. TA-1537, responds to a mutagen in nonactivation tests it will generally do so in activation tests. (The converse of this relationship is not expected.) While similar response patterns are not required for all mutagens, they can be used to enhance the reliability of an evaluation decision.

#### 4. Reproducibility

If a chemical produces a response in a single test that cannot be reproduced in one or more additional runs, the initial positive test data loses significance.

The preceding criteria are not absolute and other extenuating factors may enter into a final evaluation decision. However, these criteria are applied to the majority of situations and are presented to aid those individuals not familiar with this procedure. As the data base is increased, the criteria for evaluation can be more firmly established.



### VII. <u>EXPLANATION OF EVALUATION PROCEDURES FOR SUSPENSION ASSAYS</u>

Data obtained from mutagenicity tests are evaluated on a test by test basis followed by an examination of the total response pattern using all the data. To facilitate this type of evaluation, we have prepared two separate formats in which data are processed. The first is the Compound Summary Backup Detail Sheet, which details the essential raw data from each experiment showing surviving population counts, total mutant or convertant counts, as well as, calculated mutation frequencies. This format permits close examination of each set of test data. The following considerations are part of any assessment.

#### A. <u>Surviving Population Counts</u>

A certain level of chemically-induced toxicity is anticipated, but occasionally isolated tests or groups of tests show very low (<25%) survival compared to the tissue controls. Such isolated decreases may result from improper dilution procedures or defective growth media and decrease confidence in the calculated mutation frequencies especially if the total mutant counts appear unaffected. Data of this type are generally unacceptable and these experiments are routinely repeated at a lower dose level to reduce killing and increase confidence in the nature of the response.

#### B. <u>Total Mutant Counts</u>

For nonmutagens, the mutant/surviving population ratio should be roughly equivalent for each test point in a given experiment. If the cell number drops in response to killing, the mutant number should decrease proportionately. A mutagenic chemical, however, will produce an altered mutant/surviving population ratio. Mutant numbers as well as calculated frequencies are compared to the negative control data. In certain instances, the mutant frequencies will increase with little or no change in the absolute number of mutants especially where the test chemical is toxic. Data of this type, although not necessarily aberrant, or even rare, must be viewed with special care to ensure that the increased frequencies were not the result of selective toxicity of the test chemical for the  $\underline{\text{his}}^{\text{-}}$  cells. This phenomenon, referred to as selection, can lead to erroneous conclusions. Thus we attempt to keep the surviving population of cells high and look for positive responses that show increases in both numbers of mutants and mutation frequencies. Again, occasional isolated fluctuations in mutant counts are found that can be attributed to improper pipetting or media contamination. These fluctuations are usually easy to identify by inspection of the other data points in the experiment which will be negative.



#### C. Dose Response Phenomena

Dose-related increases in mutants and mutation frequencies are the most convincing data to have in assessing mutagenic activity of chemicals. In some cases, however, dose-related increases are not observed for mutagens. This depends considerably on the dose levels selected. The figure on the following page illustrates how one might obtain various types of dose-related responses by a mutagen based solely on dose selection. It also emphasizes the need to keep dose levels within a relatively low range of toxicity so that data are consistently on the uphill side of the hypothetical curve.

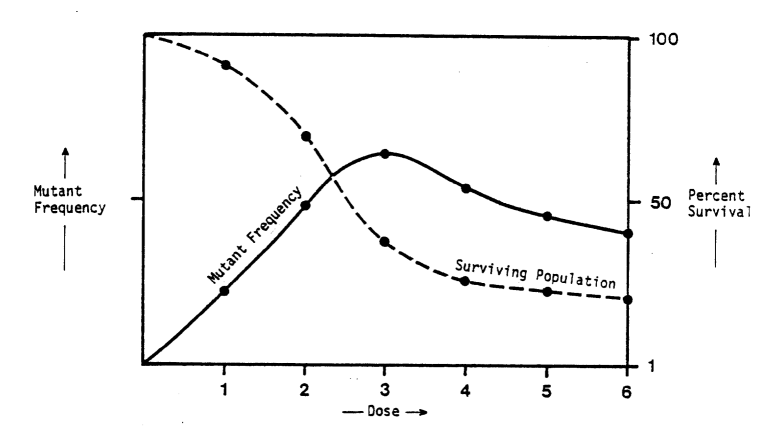
#### D. Control Tests

Positive and negative control tests are conducted with each experiment and consist of direct acting positive agents for nonactivation assays and chemicals that require metabolic transformation for activation assays. nonactivation assays, the NAN control contain the test chemical solvent plus cells, but no chemical, and is used as a reference to assess the level of response obtained in the various tests. It is not possible at this time to put precise cut-off points where negative responses become positive responses. A statistical component for our computer program is under development and will be included when available. Positive controls are only used as relative reference points and to demonstrate that the system is functioning with known mutagens. In activation assays, three types of negative controls are run: (1) A solvent control minus the chemical and minus the activation system (A-C); (2) a control plus the positive control chemical minus the activation system (A+C); and (3) a control containing the activation system and the test chemical solvent (ALI or ALU). All three controls are used collectively to assess the level of response in the various activation tests. A chemical may appear positive when compared to an A-C control but not when compared to an A+T control. The value of each of the above controls with respect to their weight in evaluation is ALI or ALU > A-C > A+C.

The other data format is the Compound Frequency Summary Report sheet in which all the calculated frequencies obtained for a given compound are displayed in a table. This format permits an overview of all data. The points form a matrix of information that should present a consistent pattern. Nonmutagens should produce a matrix with data frequencies clustered around the negative control values. Occasional random high or low fluctuations are not uncommon and seldom indicate true genetic activity. Mutagenic chemicals should, on the other hand, produce a set of consistent responses that demonstrate a logical pattern. The patterns depend on the mutagenic specificity of the chemical but can be easily recognized in the Compound Frequency Summary Report format.

These mutagenicity assays are designed to optimize the probability of recognizing mutagens from nonmutagens and, in most cases, they work well. Occasionally, the data points are such that a definitive conclusion cannot be made without additional data.





# HYPOTHETICAL EXPERIMENT

- (1) Dose levels
  1,2 & 3 were used
- (2) Dose levels
  2, 3 & 4 were used
- (3) Dose levels
  3, 4 & 5 were used

#### OBSERVED DOSE RESPONSE

A typical positive dose response set of data would be obtained.

The intermediate dose level shows a higher mutation frequency than both the low dose and the high dose.

Here an inverted dose response would be observed with the highest dose level showing the lowest response.

### APPENDIX

Tabulation of Data



# REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

			223-76-2102			PROJECT	2672	
EXPERIMEN	T 7102	03	DETECTOR TA100	SPE	CIES	/		DATE - 07/22/77
		ORG		POPU	MUTI	FRE	Q1	
COMPOUND	TEST	10	CONCENTRATION	EP+6	EP+0	EP-	8	CONTAM
	NAN		SOLVENT	0252	0216	85.	71	0
	NAP		EMS 0.066%	0616	5547	900.	49	0
000098920	NAI		0016-1 PCT.	0594	0484	81.	48	0
000098920	NA2		0008-1 PCT.	0706	0480	67.	99	0
000098920	EAN.		0004-1 PCT.	0710	0484	68.	17	0

# REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

CONTRACT EXPERIMENT 710201		T 223-76-2102 DETECTOR TA1535	SPECIES	PROJECT 2672	DATE - 07/22/77
COMPOUND	ORG TEST ID	CONCENTRATION	POPU MUTI EP+6 EP+0	• • • • • • • • • • • • • • • • • • • •	CONTAM
	NAN	SOLVENT	0810 0029	3,58	0
	NAP	EMS 0.2%	0852 5840	685.45	0
000098920	NA1	0016-1 PCT.	0924 0042	4.55	0
000098920	SAN	0008-1 PCT.	0927 0036	3.88	0
000098920	EAN	0004-1 PCT.	1036 0039	3.76	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

	CON	TRACT	223-76-2102			PROJECT	2672	
EXPERIMENT 710101		01	DETECTOR TA1537	SPECIES		/		DATE - 07/22/77
		ORG		POPU	HUTL	FRE	Qì	
COMPOUND	TEST	10	CONCENTRATION	EP+6	EP+0	EP-	8 -	CONTAM
	NAN		SOLVENT	0466	0054	11.	59	0
	NAP		QM 13 UG/ML	0235	0553	235.	32	0
000098920	NAI		0016-1 PCT.	1492	0050	3.	35	O
000098920	NAZ		0008-1 PCT.	1182	0033	2.	79	Q
000098920	EAN		0004-1 PCT.	1196	0047	3.	93	0

REPORT EXR33 LITTON BIONÈTICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT	CONTRAC 705306	T 223-76-2102 DETECTOR TA1538	SPECIES	PROJECT 2672	DATE - 07/22/77
COMPOUND	TEST ID	CONCENTRATION	POPU MUT1 EP+6 EP+0	FREQ1 EP-8	CONTAM
	NAN	SOLVENT	0403 0022	5.46	0
	NAP	NF 667 UG/ML	0376 0614	163.30	0
000098920	NAI	0016-1 PCT.	0571 0025	4.38	. 0
000098920	NA2	0008-1 PCT.	0437 0017	3.89	0
000098920	EAN	0004-1 PCT.	0449 0012	2.67	0

HEPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

CONTRACT EXPERIMENT 710202		223-76-2102 DETECTOR TA98	SPECIES	PROJECT 2672	DATE - 07/22/77
COMPOUND	OKG TEST ID	CONCENTRATION	POPU MUTI EP+6 EP+0	FREQ1 EP-8	CONTAM
	NAN	SOLVENT	0962 0133	13.83	0
	NAP	NF 667 UG/ML	0834 0598	71.70	0
000098920	NAI	0016-1 PCT.	1628 0089	5.47	0
000098920	NA2	0008-1 PCT.	1294 0134	10.36	0
000098920	EAN	0004-1 PCT.	1460 0143	9.79	O

REPORT EXR33 LITION BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

5 u o 5 o 5 u 5 u			223-76							
EXPERIMEN	1 7109	710902		DETECTOR 000004		SPECIES		/		DATE - 07/22/77
		ORG			POPU	HUTI	MUT2	FREQI	FREQ2	
COMPOUND	TEST	10	CONCEN	TRATION	EP+4	EP+1	EP+1	EP-5	EP-5	CONTAH
	NAN	•	SOLVEN	T	1175	0234	0081	19.91	6.89+	1
	NAP		EMS 1.	0 %	1463	1600	1142	109.36	78.06	0
000098920	NAI		0088-2	PCT.	1450	0582	0093	40.14	6.41	0
000098920	NA2		0044-2	PCT.	1600	0306	0090	19.13	5.63	0
000098920	NA3		0022-2	PCT.	1299	0363	0155	27.94	11.93	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT			223-76-2102 DETECTOR TA100	SPE	cies ic	DATE - 07/22/7	
COMPOUND	TEST	ORG 1D	CONCENTRATION	POPU EP+6	MUT1 EP+0	FREG1 EP-8	CONTAN
	A+C		DMN 90 UM/ML	0989	0835	84.43	0
	A-C		SOLVENT	0899	0502	55.84	0
	ALI		TISSUE	1545	0887	57.41	0
	ALU		TISSUE	1869	1097	58.69	0
	ACP	LI	DMN 90 UM/HL	1160	2112	182.07	0
	ACP	LU	DHN 90 UM/ML	0496	0456	91.94	0
000098920	ACT	LII	0016-1 PCT.	0651	0390	59.91	0
000094920	ACT	LI2	0008-1 PCT.	0540	0394	72.96	0
000098920	ACT	L13	0004-1 PCT.	0573	0407	71.03	
000098920	ACT	FAI	0016-1 PCT.	0432	0378	87.50	0
000098920	ACT	LU2	0008-1 PCT.	1450	0344	23.72	0
000098920	ACT	LU3	0004-1 PCT.	1062	0471	44.35	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT			223-76-2102 DETECTOR TA1535	SPE	CIES IC	DATE - 07/22/77	
COMPOUND	TEST	1D ORG	CONCENTRATION	POPU EP+6	MUT1 EP+0	FREQ1 EP-8	CONTAM
	A+C		DMN 90 UM/HL	0656	0025	3.81	0
	A-C		SOLVENT	0430	0025	5.81	0
	AL I		TISSUE	0617	0031	5.02	0
	ALU		TISSUE	0645	0033	5.12	0
	ACP	LI	DMN 90 UM/ML	0490	0796	162.45	0
	ACP	LU	DHN 90 UM/HL	0586	0046	7.85	0
000098920	ACT	LII	0016-1 PCT.	0571	0034	5.95	0
000098920	ACT	LIS	0008-1 PCT.	0562	0022	3.91	o
000098920	ACT	LI3	0004-1 PCT.	0706	0042	5.95	0
000098920	ACT	LUI	0016-1 PCT.	0696	0062	8.91	0
000098920	ACT	LU2	0008-1 PCT.	0559	0023	4.11	0
000098920	ACT	LU3	0004-1 PCT.	0504	0014	2.78	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

CONTRAC EXPERIMENT 710802			223-76-2102 DETECTOR TA1537	SPE	CIES ICF	PROJECT 2672 RFLO/MOUSE	DATE - 07/22/7
COMPOUND	TEST	ID OHG	CONCENTRATION	POPU EP+6	MUT1 EP+0	FREQ1 EP+8	CONTAM
	A+C		AMQ 333 UG/HL	2235	0208	9.31	o
	A-C		SOLVENT	2639	0166	6.29	0
	ALI		TISSUE	0769	0098	12.74	0
	ALU		TISSUE	0768	0054	7.03	0
	ACP	LI	AMQ 333 UG/ML	2337	2282	97.65	0
	ACP	LU	AMQ 333 UG/ML	0382	0016	4.19	2
000098920	ACT	LII	0016-1 PCT.	1561	0124	7.94	0
000098920	ACT	L15	0008-1 PCT.	0713	0118	16.55	0
000098920	ACT	LI3	0004-1 PCT.	0779	0131	16.82	0
000098920	ACT	LUI	0016-1 PCT.	1408	0151	10.72	0
000098920	ACT	LU2	0008-1 PCT.	0768	0145	18.88	0
000098920	ACT	LU3	0004-1 PCT.	0902	0169	18.74	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT		TRACT	223-76-2102 DETECTOR TA1538	SPE	CIES	PROJECT 2672 ICRFLO/MOUSE	DATE - 07/22/77
COMPOUND	TEST	ORG ID	CONCENTRATION	POPU EP+6	MUTI EP+0		CONTAM
	A+C		ANTH 67 UG/ML	0171	0026	15.20	. 0
	A-C		SOLVENT	0323	0017	5.26	0
	AL I		TISSUE	0385	0020	5.19	0
	ALU		TISSUE	0265	0031	11.70	o
	ACP	LI	ANTH 67 UG/ML	0329	0970	294.83	0
	ACP	LU	ANTH 67 UG/ML	0310	0106	34.19	0
000098920	ACT	LII	0016-1 PCT.	0361	0012	3.32	0
000098920	ACT	LIZ	0008-1 PCT.	0388	0016	4.12	o
000098920	ACT	Lla	0004-1 PCT.	0387	0014	3.62	0
000098920	ACT	LU1	0016-1 PCT.	0228	0027	11.84	0
000098920	ACT	Fn5	0008-1 PCT.	0315	0022	6.98	0
000098920	ACT	LU3	0004-1 PCT.	0249	0023	9.24	0

HFPORT EXH33 LITTON BIONETICS HUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

CONTRACT EXPERIMENT 710304			223-76-2102 DETECTOR TA98	SPE	CIES IC	DATE - 07/22/17	
COMPOUND	TEST	ORG ID	CONCENTRATION	POPU EP+6	MUT1 EP+0	FREQ1 EP-8	CONTAH
	A+C		ANTH 67 UG/HL	1797	0170	9.46	0
	A-C		SOLVENT	1513	0133	8.79	Q
	ALI		TISSUE	0810	0174	21.48	0
	ALU		TISSUE	1095	0161	14.70	0
	ACP	LI	ANTH 67 UG/ML	0629	0549	87.28	0
	ACP	ŁU	ANTH 67 UG/ML	1134	0846	74.60	0
000098920	ACT	LII	0016-1 PCT.	1398	0202	14.55	0
000098920	ACT	L12	0008-1 PCT.	0780	0164	21.03	. 0
000098920	ACT	L13	0004-1 PCT.	0777	0144	18.53	0
000098920	ACT	LUI	0016-1 PCT.	1289	0165	12.80	0
000098920	ACT	LU2	0008-1 PCT.	1501	0131	8.73	0
000098920	ACT	լսյ	0004-1 PCT.	1675	0156	9.31	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT		TRACT	223-76-2102 DETECTOR 0000D4	SPF	CIES I	12	DATE - 07/22/77		
		-			••••	J.11 E.07			DA, L
		ORG		POPU	MUTI	MUT2	FREQ1	FREQZ	
COMPOUND	TEST	10	CONCENTRATION	EP+4	EP+1	EP+1	EP-5	EP-5	CONTAM
	A+C		DHN 90 UM/ML	1184	0082	0185	6.93	15.63	0
	A-C		SOLVENT	1072	0074	0243	6.90	22.67	0
	ALI		TISSUE	1474	0153	0120	10.38	8.14	0
	ALU		TISSUE	1320	0068	0198	5.15	15.00	0
	ACP	LI	DMN 90 UM/HL	1555	0652	1122	53.36	91.82	0
	ACP	LU	DHN 90 UM/ML	1061	0202	0184	19.04	17.34	0
000098920	ACT	LII	0088-2 PCT.	1783	0103	0054	5.78	3.03	0
000098920	ACT	FIS	0044-2 PCT.	1571	0140	0042	8.91	2.67	0
000098920	ACT	LI3	0022-2 PCT.	1624	0086	0024	5.30	1.48	o
000098920	ACT	LN1	0088-2 PCT.	1909	0078	0045	4.09	2.36	0
000098920	ACT	LU2	0044-2 PCT.	1741	0077	0041	4.42	2.35	0
000098920	ACT	LU3	0022-2 PCT.	1888	0066	0031	3.50	1.64	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT 711003			223-76-2102 DETECTOR TA100	SPE	CIES SP	DATE - 07/22/77	
COMPOUND	TEST	In Oue	CONCENTRATION	P0PU EP+6	MUT1 EP+0	FREQ1 EP-8	CONTAH
	A+C		DMN 90 UM/ML	0404	0084	20.79	0
	A-C		SOLVENT	0499	0093	18.64	0
	ALI		TISSUE	0230	0192	83,48	0
	ALU		TISSUE	0487	0315	64.68	0
	ACP	LI	DMN 90 UM/HL	0319	0791	247.96	0
	ACP	LU	DHN 90 UM/ML	0779	0410	52.63	0
000098920	ACT	LII	0016-1 PCT.	0274	0126	45.99	0
000098920	ACT	LI2	0008-1 PCT.	0327	0123	37.61	0
000098920	ACT	L13	0004-1 PCT.	0418	0146	34.93	0
000098920	ACT	LUI	0016-1 PCT.	0381	0291	76.38	0
000098920	ACT	LU2	0008-1 PCT.	0492	0330	67.07	. 0
000098920	ACT	LU3	0004-1 PCT.	0436	0370	84.86	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT			223-76-2102 DETECTUR TA1535	SPE	CIES SPI	PROJECT 2672 RDAW/RAT	DATE - 07/22/77
COMPOUND	TEST	1D 086	CONCENTRATION	POPU EP+6	MUT1 EP+0	FREQ1 EP-8	CONTAM
	A+C		DMN 90 UM/ML	1388	0019	1.37	0
	A-C		SOLVENT	0841	0032	3.60	
	ALI		TISSUE	1442	0038	2.64	· •
	ALU		TISSUE	1420	0025	1.76	0
	ACP	LI	DMN 90 UM/ML	1141	4694	411.39	0
	ACP	LU	DMN 90 UM/ML	1393	0018	1.29	0
00009A920	ACT	LII	0016-1 PCT.	0995	0030	3.02	o
000098920	ACT	LI2	0008-1 PCT.	1609	0038	2.36	0
000098920	ACT	L13	0004-1 PCT.	1420	0050	3.52	0
000098920	ACT	rn1	0016-1 PCT.	1400	0028	2.00	0
000098920	ACT	LU2	0008-1 PCT.	1168	0023	1.97	0
000098920	ACT	LU3	0004-1 PCT.	1068	9026	2.43	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT		TRACT	223-76-2102 DETECTOR TA1537	SPE	CIES SPR	PROJECT 2672 Daw/Rat	DATE - 07/22/77
COMPOUND	TEST	1D ORG	CONCENTRATION	POPU EP+6	MUT1 EP+0	FREQ1 EP-8	CONTAM
	A+C		AMQ 333 UG/ML	0658	0014	2.13	0
	A-C		SOLVENT	0692	0043	6.21	0
	ALI		TISSUE	0496	0036	7.26	0
	ALU		TISSUE	0687	0018	2.62	0
	ACP	LI	AMQ 333 UG/ML	0437	0508	116.25	0
	ACP	LU	AMQ 333 UG/ML	0696	0022	3.16	0
000098920	ACT	LII	0016-1 PCT.	0469	0030	6.40	0
000098920	ACT	1.12	0008-1 PCT.	0507	0026	5.13	0
000098920	ACT	L13	0004-1 PCT.	0688	0024	3.49	0
000098920	ACT	LUI	0016-1 PCT.	0577	0035	6.07	0
00009A920	ACT	LU2	0008-1 PCT.	0783	0027	3.45	0
000098920	ACT	LU3	0004-1 PCT.	0928	0034	3.66	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM
COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT	CONTRACT 712501		223-76-2102 DETECTOR TA1538	SPE	CIES SPF	DATE - 07/22/77	
COMPOUND	TEST	ORG ID	CONCENTRATION	POPU EP+6	MUT1 EP+0	FREQ1 EP-8	CONTAM
*	A+C		ANTH 67 UG/ML	0956	0101	10.56	0
	A-C		SOLVENT	1244	0063	5.06	2
	ALI		TISSUE	0536	0058	10.82	0
	ALU		TISSUE	0552	0057	10.33	0
	ACP	LI	ANTH 67 UG/ML	0977	0869	88.95	0
	ACP	LU	ANTH 67 UG/ML	0443	0709	160.05	0
000098920	ACT	LII	0016-1 PCT.	0827	0071	8.59	2
000098920	ACT	LI2	0008-1 PCT.	0688	0101	14.68	2
000098920	ACT	LI3	0004-1 PCT.	0740	0073	9.86	0
000098920	ACT	Ful	0016-1 PCT.	0470	0079	16.81	2
000098920	ACT	LU2	0008-1 PCT.	0397	0039	9.82	0
000098920	ACT	£U3	0004-1 PCT.	0484	0063	13.02	2

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

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CONTRACT EXPERIMENT 715101			223-76-2102 DETECTOR TA98	PROJECT 2672 SPECIES SPRDAW/RAT DATE - 07/						
COMPOUND	TEST	ID ORG	CONCENTRATION	POPU EP+6	MUTI EP+0	FREQ1 EP-8	CONTAN			
	A+C		ANTH 67 UG/ML	0915	0120	13.11	0			
	A-C		SOLVENT	0405	0063	15.56	0			
	AL I		TISSUE	0455	0166	36.48	0			
	ALU		TISSUE	1127	0201	17.83	0			
	ACP	LI	ANTH 67 UG/ML	0483	1435	297.10	0			
	ACP	LU	ANTH 67 UG/ML	0961	1200	124.87				
000098920	ACT	LII	0016-1 PCT.	0454	0165	36.34	0			
000098920	ACT	F13	0008-1 PCT.	0382	0196	51.31	0			
000098920	ACT	LI3	0004-1 PCT.	0506	0179	35.38	O			
000098920	ACT	FnJ	0016-1 PCT.	0595	0206	34.62	0			
000098920	ACT	LU2	0008-1 PCT.	0938	0170	18.12	0			
000098920	ACT	LU3	0004-1 PCT.	0667	0195	29.24	0			

## REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT		NTRACT		2102 R 0000D4	SPE	CIES S	PRO VWARDAW	JECT 26	72	DATE - 07/22/77
COMPOUND	TEST	ORG ID	CONCENT	RATION	POPU EP+4	MUT1 EP+1	MUT2 EP+1	FREQ1 EP-5	FREQ2 EP-5	CONTAN
	A+C		DHN 90 L	JM/ML	1051	0164	0107	15.60	10.18	0
	A-C		SOLVENT		1533	0197	0114	12.85	7.44	0
	ALI		TISSUE		1306	0196	0135	15.01	10.34	0
	ALU		TISSUE		1091	0171	0112	15.67	10.27	0
	ACP	FI	DMN 90 U	JM/HL	1173	1294	0840	110.32	71.61	0
	ACP	LU	DMN 90 U	JH/ML	1118	0194	0086	17.35	7.69	0
000098920	ACT	LII	0088-2 P	ст.	1093	0172	0120	15.74	10.98	0
000098920	ACT	F15	0044-2 P	CT.	1125	0193	0141	17.16	12.53	0
000098920	ACT	LI3	0022-2 P	ст.	1211	0185	0091	15.28	7.51	0
000098920	ACT	լսյ	0088-2 P	cr.	1381	0185	0068	13.40	4.92	0
000098920	ACT	LU2	0044-2 Р	CT.	1131	0190	0068	16.80	6.01	0
000098920	ACT	LU3	0022-2 Р	ст.	1091	0174	0109	15.95	9.99	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT 710401			223-76-2102 DETECTOR TA100	SPE	CIES RH	PROJECT 2672 IESUS/MONKEY	DATE - 07/22/77
COMPOUND	TEST	I D ORG	CONCENTRATION	POPU EP+6	MUTI EP+0	FREQ1 EP-8	CONTAM
	A+C		DMN 90 UM/ML	0839	0737	87.84	0
	A-C		SOLVENT	0863	0557	64.54	0
	AL I		TISSUE	0874	0700	80.09	0
	ALU		TISSUE	0823	0578	70.23	0
	ACP	LI	DMN 90 UH/ML	0617	1756	284.60	0
	ACP	LU	DMN 90 UM/ML	1013	0701	69.20	0
000098920	ACT	LII	0016-1 PCT.	0959	0510	53.18	0
000098920	ACT	F15	0008-1 PCT.	0872	0663	76.03	0
000098920	ACT	LI3	0004-1 PCT.	0803	0697	86.80	0
000098920	ACT	LUI	0016-1 PCT.	0883	0576	65.23	. 0
000098920	ACT	LU2	0008-1 PCT.	0801	0653	81.52	0
000098920	ACT	LU3	0004-1 PCT.	0804	0511	63.56	O

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT			223-76-2102 DETECTOR TA1535	SPE	CTES R	PROJECT 2672 RHESUS/MONKEY	DATE - 07/22/77
COMPOUND	TEST	ORG ID	CONCENTRATION	POPU EP+6	MUT1 EP+0	FREQ1 EP-8	CONTAM
	A+C		DMN 90 UM/ML	0887	0050	5.64	0
	A-C		SOLVENT	0945	0028	2.96	0
	AL I		TISSUE	0903	0050	5.54	0
	ALU		TISSUE	0846	0032	3.78	0
	ACP	LI	DMN 90 UM/ML	1160	0660	56.90	0
	ACP	LU	DMN 90 UM/ML	1028	0045	4.38	0
000098920	ACT	LII	0016-1 PCT.	0646	0036	5.57	0
000098920	ACT	LIZ	0008-1 PCT.	1266	0047	3.71	0
000098920	ACT	LI3	0004-1 PCT.	1552	0048	3.09	0
000098920	ACT	r n I	0016-1 PCT.	0527	0032	6.07	0
000098920	ACT	LU2	0008-1 PCT.	1147	0030	2.62	0
000098920	ACT	LU3	0004-1 PCT.	1387	0055	3.97	0

HEPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMEN	CO T 710	NTRACT 901	223-76-2102 DETECTOR TA1537	SPE	ECIES RE	PROJECT 2672 IESUS/MONKEY	DATE - 07/22/77
COMPOUND	TEST	08G	CONCENTRATION	P0PU EP+6	MUT1 EP+0	FREQ1 EP-8	CONTAH
	A+C		AMQ 333 UG/ML	0352	0077	21.88	0
	A-C		SOLVENT	0641	8000	1.25	0
	AL I		TISSUE	0623	0031	4.98	0
	ALU		TISSUE	1229	0090	7.32	0
	ACP	ΓI	AMQ 333 UG/ML	1134	0428	37.74	0
	ACP	LU	AMQ 333 UG/ML	1158	0158	13.64	0
000098920	ACT	LII	0016-1 PCT.	0992	0031	3.13	0
000098920	ACT	LI2	0008-1 PCT.	0795	0044	5.53	0
000098920	ACT	LI3	0004-1 PCT.	0732	0041	5.60	_
000098920	ACT	FAI	0016-1 PCT.	0930	0032	3.44	
00009A920	ACT	LU2	0008-1 PCT.	1214	0034	2.80	. 0
000098920	ACT	LU3	0004-1 PCT.	0893	0038		0
						4.26	0

REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

CONTRACT EXPERIMENT 712503		223-76-2102 DETECTOR TA1538	SPE	CIES	PROJECT 2672 RHESUS/MONKEY	DATE - 07/22/77	
COMPOUND	TEST	ORG ID	CONCENTRATION	POPU EP+6	MUT1		CONTAN
	A+C		ANTH 67 UG/HL	0700	0097	13.86	2
	A-C		SOLVENT	0689	0079	11.47	2
	ALI		TISSUE	0590	0073	12.37	2
	ALU		TISSUE	0802	0081	10.10	2
	ACP	LI	ANTH 67 UG/ML	0798	1229	154.01	2
	ACP	LU	ANTH 67 UG/ML	1946	0081	7.74	2
000098920	ACT	LII	0016-1 PCT.	0754	0122	16.18	2
00009A920	ACT	L12	0008-1 PCT.	0766	0100	13.05	2
000098920	ACT	LI3	0004-1 PCT.	1439	0100	6.95	2
000098920	ACT	LUI	0016-1 PCT.	1118	0098	8.77	2
000098920	ACT	Fn5	0008-1 PCT.	1122	0090	8.02	2
000098920	ACT	LU3	0004-1 PCT.	1043	0090	8.63	2

HEPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

EXPERIMENT			223-76-2102 DETECTOR TA98	SPE	CIES RI	PROJECT 2672 HESUS/MONKEY	DATE - 07/22/77
COMPOUND	TEST	1D ORG	CONCENTRATION	POPU EP+6	MUT1 EP+0	FREQ1 EP-8	CONTAM
	A+C		ANTH 67 UG/ML	1852	0139	7.51	0
	A-C		SOLVENT	1660	0136	8.19	0
	AL I		TISSUE	0852	0124	14.55	0
	ALU		TISSUE	1167	0122	10.45	o o
	ACP	t I	ANTH 67 UG/ML	1150	2319	201.65	0
	ACP	LU	ANTH 67 UG/ML	1789	0147	8.22	0
000098920	ACT	LII	0016-1 PCT.	1607	0195	12.13	0
000098920	ACT	FIS	0008-1 PCT.	1455	0132	9.07	0
000098920	ACT	LI3	0004-1 PCT.	0830	0147	17.71	0
000098920	ACT	LUI	0016-1 PCT.	1579	0119	7.54	0
000098920	ACT	LU2	0008-1 PCT.	1147	0128	11.16	0
000098920	ACT	Fn3	0004-1 PCT.	1390	0141	10.14	0

## REPORT EXR33 LITTON BIONETICS MUTAGENIC ACTIVITY SYSTEM COMPOUND SUMMARY BACKUP DETAIL

CONTRACT EXPERIMENT 711302			223-76-2102 DETECTOR 0000D4	SPE	CIES R	DATE - 07/22/77			
COMPOUND	TEST	ORG ID	CONCENTRATION	POPU EP+4	MUT1 EP+1	MUT2 EP+1	FREQ1 EP-5	FREQ2 EP-5	CONTAM
	A+C		DMN 90 UM/ML	1110	0244	0121	21.98	10.90	0
	A-C		SOLVENT	1343	0108	0098	8.04	7.30	0
	AL I		TISSUE	1385	0225	0086	16.25	6.21	0
	ALU		TISSUE	1268	0263	0098	20.74	7.73	0
	ACP	LI	DMN 90 UM/ML	1370	0935	0743	68.25	54.23	0
	ACP	LU	DHN 90 UM/ML	1212	0248	0089	20.46	7.34	1
000098920	ACT	LII	0088-2 PCT.	1201	0197	0128	16.40	10.66	0
000098920	ACT	L12	0044-2 PCT.	1108	0192	0142	17.33	12.82	0
000098920	ACT	L13	0022-2 PCT.	1118	0130	0046	11.63	4.11	0
000098920	ACT	FnJ	0088-2 PCT.	0978	0220	0082	22.49	8.38	4
000098920	ACT	LU2	0044-2 PCT.	1102	0065	0036	5.90	3.27	0
000098920	ACT	LU3	0022-2 PCT.	1301	0201	0094	15.45	7.23	0